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Luca Micci, Emory University
Emily S. Ryan, Emory University
Rémi Fromentin, Université de Montréal
Steven Bosinger, Emory University
Justin L. Harper, Emory University
Tianyu He, University of Pittsburgh
Sara Paganini, Emory University
Kirk Easley, Emory University
Ann Chahroudi, Emory University
Clarisse Benne, Case Western Reserve University

Only first 10 authors above; see publication for full author list.

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Interleukin-21 combined with ART reduces inflammation and viral reservoir in SIV-infected macaques

Luca Micci,1 Emily S. Ryan,1 Rémi Fromentin,2 Steven E. Bosinger,1,3 Justin L. Harper,1 Tianyu He,4 Sara Paganini,1 Kirk A. Easley,5 Ann Chahroudi,1,6 Clarisse Benne,7 Sanjeev Gumber,8,9 Colleen S. McGary,1 Kenneth A. Rogers,1 Claire Deleage,10 Carissa Lucero,10 Siddappa N. Byrareddy,9 Cristian Apetrei,1 Jacob D. Estes,10 Jeffrey D. Lifson,10 Michael Piaatak Jr.,10 Nicolas Chomont,2 Francois Villinger,1,3 Guido Silvestri,1,3 Jason M. Brenchley,1,3 and Mirko Paiardini1,9

1Division of Microbiology and Immunology, Yerkes National Primate Research Center (YNPRC), Emory University School of Medicine, Atlanta, Georgia, USA. 2Department of Microbiology, Infectiology, and Immunology, Université de Montréal, Faculty of Medicine, and Centre de Recherche du CHUM, Montreal, Quebec, Canada. 3Yerkes Nonhuman Primate Genomics Core, Emory University, Atlanta, Georgia, USA. 4Center for Vaccine Research, University of Pittsburgh, Pittsburgh, Pennsylvania, USA. 5Department of Biostatistics and Bioinformatics, Rollins School of Public Health, and 6Department of Pediatrics, Emory University School of Medicine, Atlanta, Georgia, USA. 7Department of Pathology, Case Western Reserve University, Cleveland, Ohio, USA. 8Division of Pathology, Yerkes National Primate Research Center, Atlanta, Georgia, USA. 9Department of Pathology and Laboratory Medicine, Emory University School of Medicine, Atlanta, Georgia, USA. 10AIDS Cancer Virus Program, Frederick National Laboratory for Cancer Research, Leidos Biomedical Research Inc., Frederick, Maryland, USA.

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Despite successful control of viremia, many HIV-infected individuals given antiretroviral therapy (ART) exhibit residual inflammation, which is associated with non–AIDS-related morbidity and mortality and may contribute to virus persistence during ART. Here, we investigated the effects of IL-21 administration on both inflammation and virus persistence in ART-treated, SIV-infected rhesus macaques (RMs). Compared with SIV-infected animals only given ART, SIV-infected RMs given both ART and IL-21 showed improved restoration of intestinal Th17 and Th22 cells and a more effective reduction of immune activation in blood and intestinal mucosa, with the latter maintained through 8 months after ART interruption. Additionally, IL-21, in combination with ART, was associated with reduced levels of SIV RNA in plasma and decreased CD4+ T cell levels harboring replication-competent virus during ART. At the latest experimental time points, which were up to 8 months after ART interruption, plasma viremia and cell-associated SIV DNA levels remained substantially lower than those before ART initiation in IL-21–treated animals but not in controls. Together, these data suggest that IL-21 supplementation of ART reduces residual inflammation and virus persistence in a relevant model of lentiviral disease and warrants further investigation as a potential intervention for HIV infection.

Introduction

While combination antiretroviral therapy (ART) has dramatically improved the prognosis of HIV-infected individuals, a scalable treatment that can cure this infection is still not available. Residual disease in ART-treated HIV-infected individuals consists mainly of (a) persistent immune abnormalities, including inflammation and immune activation, limited CD4+ T cell reconstitution, and mucosal immune dysfunction (1), and (b) persistent reservoirs of latently infected cells that are not affected by ART and are responsible for the rapid rebound of virus replication upon ART discontinuation (2). Elevated levels of inflammation and immune activation that can persist even in ART-treated HIV-infected individuals with effective viral suppression have emerged as key correlates of morbidity and mortality from non-AIDS complications such as cardiovascular disease, neurocognitive dysfunction, and kidney and bone abnormalities, among others (1, 3). Furthermore, residual inflammation may contribute to HIV persistence during ART by several mechanisms, such as favoring de novo infection of activated “target” CD4+ T cells that replenish the reservoir and upregulating the expression of immune checkpoint blockers that may limit the function of HIV-specific immune responses (1, 3, 4). Therefore, a vicious cycle in which inflammation, poor antiviral responses, and HIV persistence are intimately connected may occur in ART-treated HIV-infected individuals.

IL-21 is a pleiotropic cytokine that promotes the maintenance and functionality of Th17 cells (5–7), a subset of CD4+ T cells critical for mucosal immunity (8–14). Previously, we have shown that loss of intestinal IL-21–producing cells in the gut of SIV-infected rhesus macaques (RMs) is associated with the severity of Th17 cell depletion (15). Thus, reduced IL-21 availability may contribute to the loss of antimicrobial mucosal immunity and the induction and/or maintenance of mucosal immune dysfunction and associated pathological chronic immune activation during HIV infection. Consistent with this model, we showed that administration of a rhesus IL-21–
IgFc fusion protein during acute SIV infection of RMs results in the preservation of intestinal Th17 cells, improved mucosal immune function, and reduced microbial translocation (16).

In the current study, we investigated the effects of exogenous IL-21 administration on residual immune activation and virus persistence in ART-treated, chronically SIV-infected RMs. Our findings, generated in a relevant primate model of lentiviral infection, indicate that IL-21 administration may have a significant beneficial effect on reducing residual inflammation and virus persistence during ART.

**Results**

**Experimental design and antiretroviral therapy.** Sixteen adult, Indian-origin RMs were infected i.v. with 300 50% tissue culture infective dose (TCID₅₀) SIVmac₂₅₉. Beginning on day 60 after infection (day 60 p.i.), all animals were treated with a 5-drug antiretroviral regimen — PMPA (tenofovir), FTC (emtricitabine), raltegravir, and ritonavir-boosted darunavir — for 7 months (as indicated by the gray shading in Figure 1). Eight RMs (IL-21-treated) also received rhesus IL-21-IgFc fusion protein at 100 μg/kg (s.c.), which is a dosage comparable to that used in human clinical trials (17). The IL-21-treated RMs received 2 cycles of 6 weekly doses — 1 at the time of ART initiation and 1 later on while on ART — plus 4 additional weekly doses at the time of ART interruption (orange arrows in Figure 1). The weekly dose regimen was selected on the basis of our previous study of IL-21 administration during acute SIV infection of RMs (16), as well as of our experience with in vivo administration of other cytokines in nonhuman primates (NHPs), in which weekly doses were better tolerated and showed a better response profile than did more frequent dosages (18–20). The remaining 8 RMs were treated only with ART and served as controls. Peripheral blood (PB), colorectal mucosa (RB), and lymph node (LN) biopsies were collected longitudinally and at necropsy (Figure 1A). Plasma viral load (log₁₀) before ART initiation was comparable between IL-21-treated (n = 7, animal ROC10 was excluded) and control (n = 8) RMs (4.98 ± 0.40 vs. 5.11 ± 0.40, respectively; P = 0.8354). Both ART and IL-21 were well tolerated, without major clinical complications throughout the study. Animal ROC10 in the IL-21-treated group was lost on day 140 p.i. due to postsurgical complications of a LN biopsy. Thus, all analyses after day 135 p.i. include 7 IL-21–treated and 1 control RM before infection (day 20), before infection with IL-21 treatment when assessed by flow cytometry (Supplemental Figure 4). Interestingly, IHC analyses of LN biopsies showed that IL-21–treated animals had significantly higher levels of B cells in the medulla when compared with controls (Supplemental Figure 5), thus suggesting that increased availability of IL-21 in the LN (as determined by IHC staining, data not shown) can impact the anatomic distribution of B cells during SIV infection.

**IL-21 increases intestinal Th17 and Th22 cell levels in ART-treated, SIV-infected RMs.** To assess the effects of IL-21 treatment on mucosal immunity, we next measured the levels of intestinal Th17 and Th22 cells, identified as CD4⁺ T cells producing IL-17 and IL-22, respectively, after brief in vitro stimulation with PMA and ionomycin. Representaive staining of IL-17 (Figure 2A) and IL-22 (Figure 2B) within the intestinal CD4⁺ T cells is shown in 1 IL-21–treated and 1 control RM before infection (day 20), before ART (day 58 p.i.), and at 3 time points when on ART. The reconstitution of intestinal Th17 (Figure 2C) and Th22 (Figure 2D) cells was very limited in the RMs treated with ART alone, while IL-21 supplementation was associated with a faster and more pronounced reconstitution of both cell types. Indeed, IL-21–treated RMs showed significantly higher frequencies of Th17 and Th22 in the RMs treated with ART alone versus ART plus IL-21 at all experimental time points. Thus, while ART was effective at suppressing virus replication and improved CD4⁺ T cell restoration, supplementation with IL-21 did not improve the reconstitution of CD4⁺ T cells in either blood or tissues.

**Effects of IL-21 supplementation on blood and LN CD8⁺ T cell and B cell levels.** We next assessed the effects of IL-21 on total CD8⁺ T cell and B cell levels as well as on their main differentiation subsets in PB and LN. The levels of blood (percentages and counts), LN (percentages), and RB (percentages) CD8⁺ T cells (Supplemental Figure 2) and their subsets (data not shown) were overall very similar in IL-21-treated and control RMs, with no significant differences between the 2 groups at any single experimental time point. Previous studies showed that IL-21 stimulates the production of cytotoxic molecules in CD8⁺ T cells in humans in vitro (21) and in acutely SIV-infected RMs in vivo (16). Hence, we sought to determine by flow cytometry the longitudinal expression of intracellular perforin, granzyme B (GrB), and T-bet in blood and LN CD8⁺ T cells. No significant differences were noted in the fraction of CD8⁺ T cells expressing any of these markers (Supplemental Figure 3), or a combination of them (data not shown), between IL-21–treated and control RMs. Similar to CD8⁺ T cells, the percentages of blood and LN B cells (CD20⁺), as well as their naive (CD21²CD27²), resting memory (CD21⁺CD27⁻), activated memory (CD21⁺CD27⁺), and tissue memory (CD21⁺CD27⁻) subsets (22), were also not affected by IL-21 treatment when assessed by flow cytometry (Supplemental Figure 4). Interestingly, IHC analyses of LN biopsies showed that IL-21–treated animals had significantly higher levels of B cells in the medulla when compared with controls (Supplemental Figure 5), thus suggesting that increased availability of IL-21 in the LN (as determined by IHC staining, data not shown) can impact the anatomic distribution of B cells during SIV infection.
Figure 1. Longitudinal variations in viremia and CD4+ T cell levels in IL-21–treated and control SIV-infected RMs. (A) Schematic of the study design. Sixteen RMs were infected i.v. with 300 TCID₅₀ SIVmac₂₃₉ (day 0), and starting on day 60 p.i., treated with combination ART (PMPA, FTC, raltegravir, and ritonavir-boosted darunavir) for 7 months. Seven animals (ROc10 died on day 140 p.i. due to post-surgical [LN biopsy] complications) received 2 courses of IL-21 treatment (100 μg/kg s.c.) weekly for 6 weeks at the beginning (from days 67–105 p.i.) and at the end (from days 203–241 p.i.) of ART, as well as 4 additional administrations upon ART interruption (day 271 p.i.). The remaining 8 animals served as ART-treated controls. On day 270 p.i., ART was interrupted, and all the animals were monitored for 8 additional months. PB, RB, and LN biopsies were collected at the indicated time points. (B and C) Plasma SIVmac₂₃₉ RNA levels expressed as copies/ml (LOD, 60 copies/ml, dashed line) are shown for each individual animal (B) and as an average (C) in IL-21–treated (orange circles) versus control (black squares) RMs. (D–G) CD4+ T cell levels, expressed as a fraction of live CD3+ T cells, were compared between IL-21–treated and control RMs in PB (D), RB (F), and LN (G). In PB, CD4+ T cells were also expressed as absolute counts (cells/μl blood; E). Gray shaded area represents the time of ART treatment, and the orange arrows mark IL-21 administrations. Averaged data are presented as the mean ± SEM. cART, combination ART.
Th17 and Th22 cells is a key mechanism of mucosal immune dysfunction and chronic immune activation during pathogenic HIV/SIV infections (8, 13, 14, 23, 24). To investigate whether the higher IL-21-induced numbers of intestinal Th17 and Th22 cells were associated with reductions in T cell activation and/or proliferation when on ART, we next examined the frequencies of CD4+ and CD8+ T cells expressing the activation markers HLA-DR and CD38 and/or the proliferation marker Ki-67. We found that IL-21-treated RMs showed a more rapid—i.e., already evident at day 25 on ART (day 85 p.i.)—and more pronounced reduction of activated memory (CD28CD95+CD4+ and CD8+ T cell proliferation in rectum (P ≤ 0.01; Supplemental Figure 8, A and B) and blood (P ≤ 0.01; data not shown). Interestingly, in blood (Figure 3, B and C), but not in RB (Figure 3, H and I), the reduction of immune activation was more pronounced for the key population of CM CD4+ T cells compared with that observed in EM CD4+ T cells. This was specific for CD4+ lymphocytes, since IL-21–treated RMs have lower levels of activation in both blood (Figure 3, E and F) and RB (Figure 3, K and L) CM and EM CD8+ T cells. Finally, plasma levels of soluble inflammatory markers such as IP-10, which are associated with HIV morbidity and mortality (days 143 and 200 on ART; P = 0.0401) and C-reactive protein (CRP) (day 75 on ART; P = 0.0328), were also significantly lower in IL-21–treated RMs as compared with levels in ART-only controls (Supplemental Figure 8, C and D).

These data indicate that IL-21 supplementation in ART-treated, SIV-infected RMs is associated with a more rapid and pronounced reduction of residual immune activation levels.

IL-21 treatment upregulates the expression of genes regulating antimicrobial immunity in ART-treated, SIV-infected RMs. We further explored the effect of IL-21 supplementation on ART-treated, SIV-infected RMs by comparing the whole transcriptome, as assessed by RNA sequencing (RNA-seq), in peripheral blood mononuclear cells (PBMCs) collected before ART (day 58 p.i.) and at 2 experimental time points on ART (days 50 and 200 on ART) in our 2 groups of animals. Changes in the expression of a large number of shared genes were observed in both IL-21–treated and control samples as a result of ART-induced viral suppression. Differential regulation between the 2 groups of RMs was observed in 53 genes at day 50 and in 42 genes at day 200 on ART (Q < 0.05) (Figure 4, A and E). Specifically, and consistent with the increased levels of Th17 and Th22 cell subsets (Figure 2), numerous genes associated with antimicrobial immunity were significantly upregulated in IL-21–treated animals as compared with controls at day 50 p.i. (Figure 4B). Among these, LTF, whose protein product is lactotransferrin, a major iron-binding protein with a broad range of antimicrobial and anti-inflammatory functions (25), showed a 29-fold upregulation. Of note, lactotransferrin plays a key role in immunity against Candida albicans, an important opportunistic pathogen and cause of morbidity and mortality in HIV-infected subjects (26) and against which Th17 cells are highly specialized (27).

LCN2 (lipsocalin 2) and DEFA1 (defensin α1), both of which participate in neutrophil-mediated antimicrobial immunity, were comparable to their preinfection levels. IL-21–treated animals also had significantly higher levels of CD4+IL-17+IL-22+ T cells on days 85 and 135 p.i. when compared levels in control animals (Supplemental Figure 6). The effects of IL-21 were specific for the Th17 and Th22 subsets, as the frequencies of intestinal CD4+ IFN-γ+ (Figure 2E) and CD4+IL-22+ (Figure 2F) T cells were indistinguishable between the 2 groups and were more pronounced when IL-21 was administered earlier during ART (first cycle of 6 doses), as compared with the frequencies detected during late ART (second cycle). During SIV infection of RMs, polymorphonuclear neutrophil (PMN) activation leads to accumulation of myeloperoxidase-positive (MPO+) PMNs adjacent to epithelial lesions, with the extent of MPO expression strongly correlating with the degree of intestinal epithelial barrier dysfunction (10). To assess the impact of IL-21 supplementation on this marker of mucosal immune dysfunction, we performed IHC to measure PMN infiltration in colorectal biopsy tissues from IL-21–treated and control animals before ART (day 58 p.i.) and at early (day 85 p.i.) and late (day 256 p.i.) experimental time points on ART (Figure 2G). In IL-21–treated RMs, but not in controls, the levels of MPO+ PMNs in the lamina propria (LP), in both early and late ART, were significantly lower than the levels observed on day 58 p.i. (P = 0.0078 and P = 0.0469, respectively; Figure 2H) and were thus consistent with a decrease of intestinal inflammation and, possibly, improved mucosal integrity following IL-21 treatment. Furthermore, during early ART, MPO+ PMN levels were significantly lower in IL-21–treated RMs as compared with levels in controls (P = 0.0207; Figure 2H), consistent with the higher numbers of Th17 and Th22 cells at the same experimental point. At the later time point (day 256 p.i.), ART alone became more effective at reducing MPO expression in PMNs, resulting in comparable MPO levels in the 2 groups (P = 0.6828; Figure 2H), which is consistent with the reduced differences in Th17 and Th22 cell levels at the same experimental time point.

Taken together, these data indicate that IL-21 administration improves the reconstitution of intestinal Th17 and Th22 cells and reduced intestinal inflammation in ART-treated, SIV-infected RMs.

IL-21 treatment reduces residual T cell activation and proliferation in ART-treated, SIV-infected RMs. Preferential loss of intestinal
IL-21 treatment reduces viral persistence in ART-treated, SIV-infected RMs. Recent studies have linked residual immune activation to HIV persistence (33–35). Thus, we sought to determine whether the IL-21–induced reduction of immune activation and inflammation impacted the level of virus persistence during ART by measuring plasma levels of SIV RNA using an ultrasensitive assay (LOD, 3 copies of SIV RNA/ml). As shown in Figure 5, we found that IL-21–treated RMs showed a progressive reduction in plasma SIV RNA levels compared with ART–only controls, with the fraction of RMs with an undetectable viral load remaining stable between days 75 and 200 on ART in controls (3 of 8 animals; 37.5%) but gradually increasing from 28.5% to 85.7% of animals in the IL-21–treated group over time ($P = 0.03$, test for interaction between time on study and treatment group; Figure 5A). This trend resulted in the levels of residual viremia being significantly lower in IL-21–treated RMs compared with controls by the end of the 200 days on ART ($P = 0.0203$). We next measured the frequency of cells harboring SIV DNA in the colorectum at 2 experimental time points on ART. Consistent with the data obtained for residual viremia, the reduction of SIV DNA levels (copies per 10⁴ cell equivalents) in RB biopsies between days 50 and 200 on ART was more pronounced in IL-21–treated RMs ($P = 0.0156$) than in controls ($P = 0.1484$) (Figure 5B) and therefore resulted in lower levels of SIV DNA in treated animals than in controls at day 200 on ART ($P = 0.0607$). The possibility of a link between immune activation and SIV persistence is supported by the observation that the frequencies of intestinal CD8⁺ HLA-DR⁺CD38⁺ T cells at day 50 ($r = 0.6368$; $P = 0.0107$) and day 200 ($r = 0.6314$; $P = 0.0116$) on ART positively correlated with SIV DNA content at day 200 on ART (Figure 5C).

In addition, the frequency of CD4⁺ T cells expressing PD1, a subset of cells enriched in HIV reservoirs (35), was also significantly lower in RB biopsies of IL-21–treated RMs than in controls at day 200 on ART (Supplemental Figure 10A). Of note, plasma viremia at ART initiation correlated significantly with SIV DNA contents in blood CD4⁺ T cells ($r = 0.7821$; $P = 0.0009$) and trended toward significance with SIV DNA copies in RB ($r = 0.5071$; $P = 0.0562$) at day 200 on ART, but did not correlate with the frequencies of intestinal CD8⁺ HLA-DR⁺CD38⁺ T cells at day 50 or day 200 on ART (Supplemental Table 3).

Finally, we measured the levels of CD4⁺ T cells harboring replication-competent virus using a modified quantitative viral outgrowth assay (QVOA) previously described for SIV-infected RMs (36, 37). Highly purified LN CD4⁺ T cells were isolated from 4 IL-21–treated and 4 control RMs at the last experimental time point on ART (QVOA requires a large number of cells, thus we were limited to doing this assay only in animals with a large number of cryopreserved cells). Purified CD4⁺ T cells were cocultured with the CEMx174 cell line, and frequencies of cells expressing p27 (see Figure 6A showing a representative staining), as well as levels of SIV RNA released in the supernatant, were quantified on days 9, 16, and 25 after coculture. Remarkably, all 4 IL-21–treated RMs showed levels of p27 expression and SIV RNA in supernatant that were significantly lower than those in controls at all tested experimental time points (Figure 6, B and C). As a result, the frequencies of latently infected cells, expressed as infectious units per million (IUPM) CD4⁺ T cells, were significantly lower in IL-21–treated RMs than in controls (Figure 6D).
IL-21 supplementation upregulates the expression of genes regulating antimicrobial immunity in PBMCs from ART-treated, SIV-infected RMs. (A and E) Global heatmaps of genes whose expression levels were significantly different (q < 0.05) between IL-21-treated (n = 7) and control (n = 8) RMs at day 50 (A) or day 200 (E) on ART. The specific genes are indicated on the left side of the heatmaps. (B and F) FPKM values for the selected genes of interest were compared between IL-21-treated (orange circles) and control (black squares) RMs at day 50 (B) and day 200 (F) on ART. (C and G) Gene ranking for IL-21–, IL-17–, and IL-22–related genes at day 50 (C) and day 200 (G) on ART. Dots represent individual genes plotted according to their ranking (y axis) and their addition to the cumulative score (x axis). P values below 0.25 were considered significant according to CSEA guidelines. Leading genes with higher enrichment scores in IL-21-treated RMs (as compared with controls) are highlighted in red and shown as heatmaps in B and H. Statistical analyses were performed with the Partek Genomics Suite, version 6.6, and the CSEA tool.

Along with those shown in Figure 3, these results indicate that IL-21 supplementation of ART limits both inflammation and viral persistence in SIV-infected RMs and support the possibility of a direct link between mucosal immunity, inflammation, and HIV persistence. Supporting this molecular link, SIV DNA content in purified blood CD4+ T cells at the latest time point on ART (day 256 p.i.) correlated (a) negatively with the levels of intestinal Th17 (r = −0.7632; P = 0.0013; Supplemental Figure 11A) and Th22 cells (r = −0.6429; P = 0.0116; Supplemental Figure 11C) before ART (day 58 p.i.); (b) negatively with the levels of intestinal Th17 cells on ART (day 105 p.i.; r = −0.5898; P = 0.0223; Supplemental Figure 11B); and (c) positively with the levels of intestinal CD4+Ki-67+ T cells on ART (day 84 p.i.; r = 0.5893; P = 0.0232; Supplemental Figure 11D). Moreover, at the latest time point on ART, the levels of activated (HLA-DR+CD38+) circulating CD4+ (r = 0.6885; P = 0.0062; Supplemental Figure 11E) and CD8+ (r = 0.5494; P = 0.0339; Supplemental Figure 11F) T cells positively correlated with residual plasma viremia. Finally, plasma levels of IP-10 (day 203 p.i.; r = 0.7575; P = 0.0011; Supplemental Figure 11G) and CRP (day 135 p.i.; r = 0.6986; P = 0.0050; Supplemental Figure 11H) on ART positively correlated with SIV DNA content in intestinal tissues and residual plasma viremia, respectively. Of note, plasma viremia before ART (day 58 p.i.) negatively correlated with pre-ART levels of intestinal Th17 (r = −0.6988; P = 0.0047) and Th22 (r = −0.4500; P = 0.0944) cells, but did not correlate with markers of immune activation on ART (Supplemental Table 3).

IL-21-treated RMs show better control of virus replication and immune activation after ART interruption. The design of the current study included a structured interruption of ART after 7 months as a tool to determine whether the reduced inflammation and virus persistence observed during ART in IL-21-treated RMs (Figures 3 and 5) would translate into superior control of viral replication and immune activation following ART interruption. In this part of the study, all SIV-infected RMs were monitored for 8 months after interruption of ART (off ART) and then euthanized (Figure 1). During the off-ART follow-up period, plasma viral loads were, on average, 0.7 logs lower in IL-21-treated RMs when compared with those in ART-only controls, although, due to animal-to-animal variability in this relatively limited cohort, the difference between the 2 groups did not reach statistical significance at any of the experimental time points (Figure 7A). The difference was particularly pronounced starting at day 60 off ART, after which viral loads consistently increased (up to 1 log) in controls but remained stable in IL-21-treated animals (Figure 7A). To further investigate potential differences in the viral rebound after ART interruption, we then compared the levels of plasma viremia at different experimental time points off ART with the pre-ART levels. We found that plasma viremia was significantly reduced as compared with pre-ART levels up to day 60 off ART in both groups (data not shown); however, starting at day 90 off ART, plasma viremia rebounded to levels similar to those observed before ART in control RMs, while IL-21-treated animals maintained plasma SIV RNA copies that were significantly lower than the pre-ART baseline up to day 240 off ART (Figure 7B). As expected, plasma viremia levels before ART (day 58 p.i.) strongly correlated with those after ART interruption (shown for days 90 and 240 off ART in Supplemental Figure 12), thus highlighting the critical importance of matching experimental groups for viremia before ART initiation when testing interventions, as in our study. We next measured the levels of SIV DNA in purified blood CD4+ T cells after ART interruption (Figure 7C) and found that, in the majority of control RMs, these levels increased as compared with their on-ART levels (P = 0.0031; day 180 off ART), and, in fact, they were no longer different from pre-ART levels (P = 0.0829). In contrast, in IL-21-treated RMs, the levels of SIV DNA in blood CD4+ T cells did not increase following ART interruption (P = 0.1931 vs. on-ART levels) and remained significantly lower (P = 0.0407) compared with pre-ART levels (Figure 7C). Interestingly, in 2 control animals, the levels of CD4+ T cell SIV DNA content decreased after ART interruption. Both animals were Mamu-A*01+ and had among the lowest set-point viral loads before ART initiation, thus suggesting the possibility of a genetically predisposed control of viral replication (38). Of note, when considering only Mamu-A*01+ animals, we found that the levels of SIV DNA content off ART rebounded to levels significantly higher than on-ART levels in all 4 controls, but only in 1 of the 4 IL-21-treated animals (Figure 7D). As a result, CD4+ T cell–associated SIV DNA content on day 180 after ART interruption was significantly lower in IL-21-treated animals compared with control RMs (mean ± SEM log_{10}: 3.11 ± 0.52 vs. 4.55 ± 0.11; P = 0.01; Figure 7D).

We next measured, in the same SIV-infected RMs, the levels of activated or proliferating T cells after ART interruption. We found that IL-21-treated animals maintained CD4+HLA-DR+CD38+ and CD8+HLA-DR+CD38+ T cell levels that were significantly lower than those observed in controls in both PB and RB and up to day 240 off ART, i.e., 7 months after the last dose of IL-21 (Figure 7E). We also found that the fraction of CD4+HLA-DR+CD38+ T cells at necropsy was significantly lower in the LNIs of IL-21-treated RMs compared with that detected in control RMs (P = 0.0260; Supplemental Figure 13A), with a similar trend found for CD8+HLA-DR+CD38+ T cells (Supplemental Figure 13B). Finally, the improved control of chronic immune activation and viral replication resulted in a higher CD4+/CD8+ ratio after ART interruption in IL-21-treated RMs, with values that become statistically significant at day 180 off ART compared with control RMs (Supplemental Figure 13C). Together, these data suggest that IL-21 supplementation of ART results in beneficial effects upon ART discontinuation, in particular in reducing chronic immune activation.
SIV-infected RMs and determined how this intervention impacted immune reconstitution, residual immune activation, and viral persistence during ART as well as after ART interruption. The rationale for using IL-21 was 4-fold. First, this cytokine regulates processes that are compromised during pathogenic HIV/SIV infections, including differentiation of Th17 cells (5–7), maintenance of functional CD8+ T cells (39–42), and differentiation of memory B cells and Ab-secreting plasma cells (43–46). Second, SIV infection is associated with a loss of IL-21-producing cells in the gut, which is correlated with the severity of intestinal Th17 cell depletion (15). Third, administration of IL-21 during acute SIV infection of RMs is discussed.

Discussion

While ART has a major beneficial impact on HIV disease progression, this treatment is associated with residual morbidity related to a number of persistent immune abnormalities and to the presence of a reservoir of latently infected cells that will cause a rebound of virus replication if ART is interrupted. As such, numerous concepts and products are aggressively being explored as novel interventions to be added to standard ART in order to reduce or possibly eliminate these immunological and/or virological aspects of residual HIV disease on ART. In the current study, we administered IL-21, a potent immunomodulatory cytokine, to ART-treated, SIV-infected RMs and determined how this intervention impacted immune reconstitution, residual immune activation, and viral persistence during ART as well as after ART interruption. The rationale for using IL-21 was 4-fold. First, this cytokine regulates processes that are compromised during pathogenic HIV/SIV infections, including differentiation of Th17 cells (5–7), maintenance of functional CD8+ T cells (39–42), and differentiation of memory B cells and Ab-secreting plasma cells (43–46). Second, SIV infection is associated with a loss of IL-21-producing cells in the gut, which is correlated with the severity of intestinal Th17 cell depletion (15). Third, administration of IL-21 during acute SIV infection of RMs is
associated with preserved intestinal Th17 cells and limited microbial translocation (16). Last, IL-21 is currently being evaluated in several phase I and II cancer clinical trials, with limited toxicity and encouraging single-agent activity (17, 47–49). Importantly, a historic limitation of the SIV/RM model for studies of HIV residual disease, i.e., the lack of an optimized ART that fully suppresses virus replication, has been recently overcome, with a number of studies showing that the combination of tenofovir, emtricitabine, an integrase inhibitor (raltegravir or dolutegravir), and darunavir can durably suppress SIV infection of RMs to clinically relevant levels (50–53). Indeed, in the current study, we achieved prolonged suppression of plasma viral replication (<60 SIV RNA copies/ml) in all SIV<sub>mac</sub>-infected RMs, highlighting the relevance of the ART-treated, SIV-infected RM model to test immune-based interventions targeting residual immune activation and virus persistence in HIV-infected humans on suppressive ART.

IL-21 administration was safe in the context of ART-treated, SIV infected RMs. When compared with ART-only controls, RMs receiving ART plus IL-21 showed (a) better reconstitution of intestinal Th17 and Th22 cells and a selective enrichment in genes that participate in Th17 and Th22 cell–orchestrated antimicrobial immunity, and (b) a significantly more rapid and pronounced reduction of the levels of cellular immune activation in rectum and blood, as well as of plasma levels of the inflammatory markers associated with preserved intestinal Th17 cells and limited microbial translocation (16). Last, IL-21 is currently being evaluated in several phase I and II cancer clinical trials, with limited toxicity and encouraging single-agent activity (17, 47–49). Importantly, a historic limitation of the SIV/RM model for studies of HIV residual disease, i.e., the lack of an optimized ART that fully suppresses virus replication, has been recently overcome, with a number of studies showing that the combination of tenofovir, emtricitabine, an integrase inhibitor (raltegravir or dolutegravir), and darunavir can durably suppress SIV infection of RMs to clinically relevant levels (50–53). Indeed, in the current study, we achieved prolonged suppression of plasma viral replication (<60 SIV RNA copies/ml) in all SIV<sub>mac</sub>-infected RMs, highlighting the relevance of the ART-treated, SIV-infected RM model to test immune-based interventions targeting residual immune activation and virus persistence in HIV-infected humans on suppressive ART.

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residual immune activation and virus persistence during ART. While, on the basis of our data, we propose increased mucosal integrity as a key contributor to the reduced residual immune activation during ART in IL-21-treated RMs, the mechanisms by which a reduction of immune activation can limit virus persistence under ART may be very complex and may include (a) a reduction in the number of activated CD4+ T cells that serve as targets for the infection and that may become a persistent reservoir, and (b) a better maintenance and/or restoration of HIV/SIV-specific immune responses, thus resulting in a better clearance of productively infected cells and/or latently infected cells in which virus infection has been reactivated. Although this study was not aimed at determining the mechanism(s) of the link between residual immune activation and virus persistence, further studies that include elective necropsy of the SIV-infected RMs during ART may be able to provide critical insights on this important question. Moreover, the activity of IL-21 in increasing mucosal integrity was more pronounced when IL-21 was administered earlier during ART as compared with administration late in ART. Thus, further studies are needed to confirm the benefit of IL-21 administration on mucosal integrity in already fully suppressed SIV-infected animals. As for all therapeutic interventions improving reconstitution of CD4+ T cells during ART, one potential concern is that, by increasing Th17 cell levels, we also expand the viral reservoir harbored by these cells. In this regard, it is important to note that Th17 cell levels are never greater than those detected before SIV infection; thus, IL-21 treatment does not induce a nonphysiological accumulation of Th17 cells.

An additional and perhaps surprising result of this study is that the beneficial effects of IL-21 supplementation seem to include better control of immune activation and, although to a lower extent, viral replication following ART interruption. This observation is intriguing in that it supports the idea that immune-based interventions during ART-induced suppression of virus replication may have an impact on the dynamics of the host-virus interaction that persists even after ART discontinuation, when virus replication has rebounded to levels similar to those observed before ART initiation. The observation that IL-21 supplementation might have caused a significant and persistent reduction in immune activation despite robust virus replication is reminiscent of the phenotype typical of the nonpathogenic SIV infection of natural hosts such as the sooty mangabeys (54). Of note, in both IL-21-treated and control RMs, plasma viremia at ART initiation directly correlated with the extent of viral rebound after ART interruption, thus highlighting the critical importance of matching experimental groups before ART initiation in studies aimed at testing the curative potential of therapeutic interventions. Future studies in which the ART plus IL-21-treated, SIV-infected RMs are followed up for longer periods after ART interruption will be needed to determine the impact of this treatment on the clinical progression of SIV infection.

In summary, we tested a novel immune-based intervention in an in vivo NHP model highly relevant for studies of HIV residual immune dysfunction and virus persistence under ART. By using the well-established model of SIV infection in RMs in the context of ART-mediated suppression of virus replication, we determined that IL-21 supplementation is associated with key benefits in terms of reducing both the residual immune activation and inflammation and the level of virus persistence in ART-treated, SIV-infected RMs. In addition, these in vivo data support the hypothesis of a mechanistic link between residual immune activation and virus persistence that may have important implications in the pathogenesis of residual HIV disease under ART. Taken together, these results provide a rationale for further exploration of IL-21 as an immune-based intervention in preclinical and clinical studies of HIV/SIV eradication.

Methods

Animals, SIV infection, and ART regimen. Sixteen RMs, all housed at the YNPRC in Atlanta, Georgia, were included in the study. All ani-
mals were *Mamu-B*‘08 and ‘B’17, while 8 of them were *Mamu-A*‘01 (RLm12, RBt12, RP11, RChb2, RV110, RKg11, ROC10, and RP8). The 16 RMs were randomized into 2 groups (group 1: IL-21 plus ART treatment; group 2: ART only) of 8 animals on the basis of age (group 1: 6.9 ± 0.63 years; group 2: 6.7 ± 0.86 years), weight (7.4 kg ± 0.6 kg vs. 6.7 kg ± 0.3 kg), and *Mamu-A*‘01 status (4 in each group). All 16 animals were infected i.v. with 300 TCID_{50} SIVmac~199~ (day 0). Starting on day 60 p.i., all animals were treated with a 5-drug ART regimen consisting of 2 reverse transcriptase (RT) inhibitors (PMPA, 20 mg/kg and FTC, 30 mg/kg), 1 integrase inhibitor (raltegravir, 100 mg/kg), and 1 protease inhibitor (darunavir, 375 mg/bid with ritonavir, 50 mg/bid, as a boosting supplementation) for 7 months. In addition, the 8 animals in group 1 received 2 cycles of treatment with recombiant IL-21–IgFc (rIL-21–IgFc) (16) (100 μg/kg s.c., once per week for 6 weeks) at the beginning (from days 67 to 105 p.i.) and at the end (from days 203 to 241 p.i.) of ART, as well as 4 additional administrations upon ART interruption (day 271 p.i. or day 1 after ART discontinuation). The remaining 8 animals in group 2 served as ART-treated controls (Figure 1A). We did not notice any side effects or increased complications in the SIV-infected RMs that had been treated with IL-21 compared with those treated with ART alone. ROC10, in the IL-21–treated group, was euthanized on day 140 p.i. due to post-surgical (LN biopsy) complications. Markers in this animal were comparable to those of the others treated with IL-21, including greater Th17 and Th22 cell levels compared with those in ART-treated controls (Supplemental Table 4). On day 270 p.i., ART was interrupted, and all animals were monitored for an additional 8 months. PB, RB, and LN biopsies were collected at numerous experimental time points throughout the study (Figure 1).

**Production and testing of rhesus rIL-21–IgFc.** Rhesus rIL-21–IgFc (IL-21) fusion protein was generated as previously described by the Resource for Nonhuman Primate Immune Reagents of Emory University (16). Briefly, IL-21 was produced in the *Drosophila* S2 system as a fusion protein between r*MamuIL*-21 and a macaque IgG2 Fc mutated to prevent binding to complement or Fc receptors, similar to a previously published report on rPD1–IgFc (55). IL-21–Fc was purified to greater than 95% by protein G sepharose affinity chromatography, dialyzed against PBS, and tested for sterility and the potential presence of residual endotoxin (16).

**Sample collection and processing.** The collection and processing of PB and LN were done as previously described (8, 15, 16, 56). Briefly, blood samples were used for a complete blood count and routine chemical analysis and plasma separated by centrifugation within 1 hour of collection. PBMCs were prepared by density gradient centrifugation. Up to 20 RB biopsies were collected with a biopsy forceps under visual control via an anoscope. RB-derived lymphocytes were isolated by digestion with 1 mg/ml collagenase for 2 hours at 37°C and then passed through a 70-μm cell strainer to remove residual tissue fragments. For LN biopsies, the skin over the axillary or inguinal region was clipped and surgically prepared. An incision was made over the LN, which was homogenized and passed through a 70-μm cell strainer to mechanically isolate lymphocytes. All samples were processed, fixed in 1% paraformaldehyde, and analyzed within 24 hours of collection.

**Flow cytometric analysis.** Fourteen-parameter flow cytometric analysis was performed on PB-1, PBMC-1, LN-, and RB-derived cells according to standard procedures using a panel of mAbs that we and others have shown to be cross-reactive with RMs (15, 16, 56, 57). Predetermined optimal concentrations of the following Abs were used: anti-CD3–APC–Cy7 (clone SP34-2), anti-CD95–PE–Cy5 (clone DX2), anti-CD28–PE–594 (clone CD28.2), anti-CCR5–APC (clone 3A9), anti-CCR5–PE (clone 3A9), anti-Ki–67–Alexa Fluor 700 (clone B56), anti–IFN-γ–PE–Cy7 (clone B27), anti–CD8–PE–CF–594 (clone RPA-T8), anti–TNF-α–Alexa Fluor 700 (clone MAb11), anti–CCR7–PE–Cy7 (clone 3D12), anti–HLA–DR–PerCp–Cy5.5 (clone G46-6) (all from BD Pharmingen); anti–IL-17–Alexa Fluor 488 (clone eBio-64DEC17), anti–IL-22–APC (clone IL220P) (all from eBioscience); anti–CD4–BV421 (clone OKT4), anti–CD4–BV605 (clone OKT4), anti–CD1–PE (clone EH12.2H7), anti–CD4–BV421 (clone EH12.2H7), anti–IL-2–BV605 (clone MQ1-1H12), anti–CD20–PerCp–Cy5.5 (clone 2H7) (all from BioLegend); anti–CD8–Qdot705 (clone 3B5) and Aqua LIVE/DEAD amine dye AmCyan (both from Invitrogen); and anti–CD38–FITC (clone AT-1; STEMCELL Technologies). Flow cytometric acquisition was performed on at least 100,000 CD3+ cells on a BD LSR II Flow Cytometer driven by BD FACSDiva software. Analysis of the acquired data was performed using FlowJo software.

**Intracellular cytokine staining.** Th17 and Th22 cell levels were determined as the percentage of CD4+ T cells that produced IL-17 and IL-22 following in vitro stimulation with PMA and ionomycin (16). PBMCs, LN-, and RB-derived cells, isolated as described above, were resuspended to 3 × 10^6 cells/ml in complete RPMI 1640 medium. Cells were then incubated for 4 hours at 37°C in medium containing PMA, A23187, and BD GolgiStop. Following incubation, the cells were washed and stained with surface markers for 30 minutes in the dark at room temperature, followed by fixation and permeabilization. After permeabilization, cells were washed and stained intracellularly with Abs against the cytokines of interest for 1 hour in the dark at room temperature. Following staining, cells were washed, fixed in PBS containing 1% paraformaldehyde, and acquired on a BD LSR II Flow Cytometer.

**Plasma levels of immune activation and inflammation markers.** Soluble IFN-γ–induced protein 10 (IP-10) and CRP levels were measured in plasma using commercially available ELISA kits according to the manufacturer’s instructions. IP-10 levels were quantified using a human IP-10 Quantikine ELISA kit (R&D Systems) and expressed as pg/ml. Plasma CRP levels were measured using a monkey CRP ELISA kit (Life Diagnostics Inc.) and expressed as μg/ml.

**Plasma viral load and tissue-associated SIV DNA/RNA in rectum.** Plasma SIV viral loads were determined by standard quantitative RT-PCR, as previously described (58). Ultrasonic presentations of plasma viral loads were achieved by concentrating virus from the larger volumes of material available by ultracentrifugation as previously described (53). Quantitative assessments of SIV DNA and SIV RNA in mucosal tissues at days 50 and 200 on ART were determined by quantitative hybrid real-time/digital RT-PCR and PCR assays, as previously described (53). For each sample, 12 replicate reactions were run with a nominal single-copy sensitivity. The clinical sensitivity (based on the number of cells assessed) in our samples was as low as 1 copy/850,000 cells.

**Cell-associated SIV DNA within blood CD4+ T cells.** Quantitative assessment of cell-associated total SIV DNA within circulating CD4+ T cells at day 58 p.i., at days 50 and 200 on ART, and at days 90 and 180 off ART was performed using a modified version of a recently published quantitative nested PCR assay for cell-associated total HIV DNA (59). In a first round of PCR, total SIV DNA was amplified with 2 primers that anneal within a conserved region of the long terminal repeat...
(LTR) 5' end (SIV-LF1) and at the junction with the Gag gene (SIV-R1). The forward primer SIV-LF1 was extended with a lambda phage-specific heel sequence at the 5' end of the oligonucleotide. Primers targeting the CD3 gene (HC3DOUT-5' and HC3DOUT-3') were also added to quantify the exact number of cells in the initial samples. Gag-LTR sequences were amplified from 15 μl lysate in a 50-μl reaction mixture composed of 1X Taq Buffer, MgCl₂, dNTP, SIV-LF1, SIV-R1, and Taq polymerase. The first-round PCR cycle conditions were as follows: a denaturation step of 8 minutes at 95°C and then 16 cycles of amplification (95°C for 1 minute, 62°C for 40 seconds, 72°C for 1 minute), followed by an elongation step at 72°C for 15 minutes. In a second round of PCR, the lambda T-specific primer (Lambda T) and the LTR primer (SIV-R2) were used to amplify SIV sequences obtained from the first amplification. Primers targeting CD3 were also used in another second-round PCR. Nested PCR was performed on one-tenth of the first-round PCR product in a mixture consisting of 1X Rotor-Gene Master Mix (QIAGEN), Lambda T primer, SIV-R2 primers, and SIV probe (Integrated DNA Technologies). For CD3 amplification, nested PCR was performed in a mixture composed of 1X Rotor-Gene Master Mix, HC3DIN 5' and MamuCD3IN 3' and MamuCD3 probe. Cycling was performed on the Rotor-Gene (QIAGEN) with a denaturation step (95°C for 4 minutes), followed by 40 cycles of amplification (95°C for 3 seconds, 60°C for 10 seconds). The total SIV DNA copy number was calculated using a standard curve as a reference. This standard curve consisted of serial dilution of the 3D8 cell lysates (carrying 1 integrated copy of SIV genome per cell) (60).

Quantification of replication-competent virus in purified CD4+ T cells during ART. LN cryopreserved, highly purified CD4+ T cells were cocultured at a 1:1 ratio with the CEMx174 cell line (NIH AIDS Research and Reference Reagent Program) in serial dilutions ranging from 1 × 10⁶ to 0.1 × 10⁶ cells per well. The cells were cultured in complete RPMI 1640 with 4 mM L-glutamine supplemented with 10% heat-inactivated FBS (Gemini Bio Products), penicillin (50 U/ml), streptomycin (50 μg/ml), and IL-2 (100 U/ml). Cultured cells were split, fed with fresh medium, and harvested and analyzed on days 9, 16 and 25. Flow cytometric analyses of intracellular SIV-Gag p27 expression (61) and SIV-Gag viral RNA (vRNA) (copies/ml) in the supernatant were used to determine positive wells. Replication-competent SIV levels in CD4+ T cells were compared between cell cultures from controls and IL-21-treated animals. The frequencies of infected cells were determined by the maximum likelihood method (62) and expressed in terms of IUPM CD4+ T cells.

RNA-seq analysis. RNA-seq analysis was conducted at the Yerkes Nonhuman Primate Genomics Core laboratory (http://www.yerkes.emory.edu/nhp_genomics_core/). PBMCs at various time points were stored in QIAGEN RLT buffer at -80°C. Total RNA was prepared using QIAseq RNaseq kits. Polyadenylated transcripts were purified on oligo-dT magnetic beads, fragmented, reverse transcribed using random hexamers, and incorporated into barcoded cDNA libraries based on the Illumina TruSeq platform. Libraries were validated by microelectrophoresis and quantified, pooled, and clustered on Illumina TruSeq v3 flow cells. Clustered flow cells were sequenced on an Illumina HiSeq 1000 in 100-base single-read reactions. RNA-seq data were analyzed by alignment and annotation to the MacaM assembly of the Indian rhesus macaque genome (63) (available at http://www.umce.edu/rhesusgenechip/index.htm#NewRhesusGenome).

Alignment was performed using STAR version 2.3.0e (64); parameters were set using the annotation as a splice junction reference, and unannotated, noncanonical splice junction mappings and nonunique mappings were removed from downstream analysis. Transcripts were annotated using MacaM assembly and annotation, version 7.6 (see previous URL address). RNA-seq reads and the normalized expression table were deposited in the NCBI’s Gene Expression Omnibus (GEO) database (GEO GSE73232). Transcript assembly, abundance estimates, and differential expression analysis were performed using Cufflinks, version 2.1.1, and Cuffdiff (65). RNA-seq data are expressed as fragments per kilobase mapped (FPKM), which is the default output by Cufflinks/Cuffdiff, in which reads and fragments mapping to an individual gene are normalized by the total number of kilobases that map to the reference genome for an individual sample; this calculation minimizes bias due to differences in sequencing depth between samples. Heatmaps and other visualizations were generated using the Partek Genomics Suite, version 6.6.

GSEA. GSEA was performed using the desktop module available from the Broad Institute (www.broadinstitute.org/gsea/). While conventional statistics for differential gene expression determine significance on the basis of variance on a single gene, GSEA uses a cumulative statistic based on multiple genes within a biological pathway, and these multiple genes are grouped together on the basis of their common function/characteristic or empirical data (66). Gene rank was calculated for the normalized expression table using the signal-to-noise metric. The ranked dataset was screened against the Broad Institute’s Molecular Signatures Database (http://www.broadinstitute.org/gsea/msigdb/index.jsp), curated (C2) and immunologic signatures (C7) gene sets, as well as against the ISG gene set, an immune activation dataset determined using previous data from our laboratory (67). We also generated unbiased sets of genes representing IL-21-regulated genes for GSEA testing. Gene sets with P values below 0.25 were considered significant according to the guidelines published on the GSEA home page.

IHC and quantitative image analysis. IHC and quantitative image analysis for the percentage of LP area that stained for MPO+ neutrophils was performed as previously described (68, 69).

Statistics. Repeated-measures analyses for each outcome (CD4+ T cells, CD4+Ki-67+ T cells, Th17 cells; plasma viral load, SIV DNA within blood CD4+ T cells, etc.) were performed with a means model using SAS PROC MIXED, version 9 (SAS Institute) providing separate estimates of the means by weeks after infection and treatment groups. A compound-symmetry variance-covariance form in repeated measurements was assumed for each outcome, and robust estimates of the standard errors of parameters were used to perform statistical tests and construct 95% CIs (70). The mean estimates and their 95% CIs were back transformed to the original scale and reported as the geometric mean with 95% CIs. The model-based means are unbiased with unbalanced and missing data, so long as the missing data are noninformative (missing at random). t tests were used to compare the differences between the model-based treatment means (least-squares means) at each time point and to compare mean differences over time within each treatment group. Specific statistical tests were generally performed within the framework of the mixed-effects linear model. However, selected statistical tests were performed for specific data subsets. All statistical tests were 2 sided and unadjusted for multiple comparisons. A P value below 0.05 indicated statistical significance. Because of the small sample sizes, results from these studies focus on the magnitude of the differences for each outcome, consistency of the findings, and biological significance. The generalized estimat-
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Study approval. This study was approved by the IACUC of the YNPRC at Emory University (permit 2001973). All animal experimentations were conducted following guidelines established by the Animal Welfare Act and the NIH for the housing and care of laboratory animals and performed in accordance with institutional regulations. All efforts were made to minimize suffering of the animals.

Author contributions

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Address correspondence to: Mirko Paiardini, Division of Microbiology and Immunology, Yerkes National Primate Research Center, Emory University School of Medicine, 954 Gatewood Road, Atlanta, Georgia 30329, USA. Phone: 404.727.9840; E-mail: mirko.paiardini@emory.edu.

Siddappa N. Byrareddy’s present address is: Department of Pharmacology and Experimental Neuroscience, University of Nebraska Medical Center, Omaha, Nebraska, USA.

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